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INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification 6:

A61K 31/44, 47/32, 9/48

(11) International Publication Number: WO 99/61022

(43) International Publication Date: 2 December 1999 (02.12.99)

(21) International Application Number: PCT/IB99/00139

(22) International Filing Date: 26 January 1999 (26.01.99)

(30) Priority Data:

09/086,224

28 May 1998 (28.05.98)

US

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(81) Designated States: AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, US, UZ, VN, YU, ZW, ARIPO patent (GH, GM, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

Published

With international search report.

(54) Title: A STABLE ORAL PHARMACEUTICAL COMPOSITION CONTAINING A SUBSTITUTED PYRIDYLSULFINYL BENZIMIDAZOLE

(57) Abstract

A pharmaceutical composition which is stable and suitable for oral administration to a patient comprises a mixture of a substituted pyridyl sulfinyl benzimidazole having gastric acid secretion inhibitory activity (such as omeprazole, lansoprazole, or pantoprazole), and a pharmaceutically acceptable carrier. The carrier comprises a polymer having vinyl pyrrolidone monomeric units, such as polyvinylpyrrolidone or a vinyl pyrrolidone–vinyl acetate copolymer. Surprisingly, it has been found that the vinylpyrrolidone polymer acts as a stabilizing excipient on the substituted pyridyl sulfinyl benzimidazole so that the composition need not include any alkaline components to prevent degradation of the active ingredient. In a preferred embodiment, the composition is in the form of a capsule, whereby the mixture of the substituted pyridyl sulfinyl benzimidazole and the vinyl pyrrolidone polymer in the form of a powder blend or granules, is contained within a capsule shell, which capsule shell is made from an enteric material or is coated with an enteric material.

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A STABLE ORAL PHARMACEUTICAL COMPOSITION CONTAINING A SUBSTITUTED PYRIDYLSULFINYL BENZIMIDAZOLE

5 Background of the Invention

The present invention relates to a stable oral pharmaceutical composition comprising a substituted pyridylsulfinyl benzimidazole having gastric acid secretion inhibitory activity as the active ingredient and a carrier which acts as a stabilizing excipient. The invention also relates to a process for making the pharmaceutical composition.

10

United States Patent Nos. 4,255,431; 4,628,098; and 4,758,579 disclose substituted pyridylsulfinyl benzimidazoles (such as omeprazole) as potent inhibitors of gastric acid secretion. This class of compounds inhibits gastric acid secretion by inhibiting H*-K* ATPase (proton pump) activity. Drugs in this class are known to be highly unstable in an acidic environment. They are also unstable in the presence of moisture and organic solvents. Thus, the formulation in which the drugs are to be administered to a patient, and the process for manufacture of the formulation, must be designed to protect the drug from moisture as well as an acidic environment. Due to the very rapid drug degradation which occurs in acidic gastric fluids, the formulations should also be enteric coated.

20

United States Patent No. 4,786,505 discloses an oral pharmaceutical composition comprising a core containing omeprazole together with an alkaline reacting compound, or an alkaline salt of omeprazole optionally together with an alkaline compound; one or more subcoating layers comprising inert reacting compounds which are soluble or rapidly disintegrating in water, or polymeric, water soluble film forming compounds, optionally containing pH-buffering alkaline

compounds; and an outer enteric coat. The alkaline reacting compound is a pharmaceutically acceptable substance (or substances) which creates a "micro-pH" around each omeprazole particle of not less than pH=7, preferably not less than pH=8, when water is adsorbed onto the particles of the mixture or when water is added in small amounts to the mixture. The subcoating layer separates the omeprazole containing core from the enteric coating polymer(s) containing free carboxyl groups. The enteric coating polymers can otherwise cause degradation of omeprazole during the coating process or during storage.

Japanese Patent 05-194,225 discloses tablets, granules and capsule formulations where the 10 benzimidazole gastric ulcer inhibitors are stabilized by compounding with amino acids and buffering agents.

United States Patent No. 5,385,739 discloses a stable microgranule formulation containing a neutral core of sugar and starch and an active layer consisting of a dilution of omeprazole in mannitol in substantially equal amounts, wherein the active omeprazole layer contains about 10% by weight of carboxymethylstarch, and about 5% by weight of sodium lauryl sulfate, and wherein the dilution of omeprazole in mannitol is applied to the neutral core by means of hydroxypropyl methylcellulose as a high viscosity binder.

20 PCT Int. Pat. Appl. WO 97/12581 discloses a composition comprising: (a) a core containing omeprazole as the active principle, the core being constituted of nuclei and the omeprazole active principle mixed together and then compressed together, the omeprazole active principle not being in the form of an alkaline salt; (b) an intermediate layer; and (c) an enteric layer. The

composition disclosed therein is stated to be free of alkaline reacting compounds which had

previously been considered as essential; however, each of the compositions exemplified in

W0 97/12581 contains either a lubricant, such as sodium stearyl fumarate, magnesium stearate, or

talc in the core, or talc in the intermediate layer. These compounds are alkali metal or alkaline

5 earth metal salts and are known to be alkaline in nature.

It is an object of the present invention to provide a stable oral pharmaceutical composition

containing a substituted pyridylsulfinyl benzimidazole having gastric acid secretion inhibitory

activity as the active ingredient and a carrier, which composition is free of alkaline compounds.

10

It is a further object of the present invention to provide a process for the preparation of a stable

oral pharmaceutical composition in the form of a mixture containing a substituted pyridylsulfinyl

benzimidazole and a carrier, wherein the mixture is not compressed to form hard intact core units,

but is filled in the form of a powder or granules into enteric-coated capsules or capsules made

15 from an enteric material. Once the capsules dissolve, the drug particles are freely dispersed in the

gastrointestinal fluid so as to result in a rapid rate of dissolution and absorption of the drug.

It is a further object of the present invention to provide a process for the preparation of a stable

oral pharmaceutical composition in the form of a mixture containing substituted pyridylsulfinyl

20 benzimidazole which process is simple, less time consuming, and more economical than prior art

processes.

Summary of the Invention

It has been surprisingly found that in a mixture comprising a substituted pyridylsulfinyl benzimidazole and one or more polymers obtained by the polymerization of monomers at least one of which is vinylpyrrolidone, the substituted pyridylsulfinyl benzimidazole is stabilized. The 5 mixture, even though it is free of alkaline reacting compounds, does not show a change in color which is typically observed in compositions where the benzimidazole has undergone degradation.

Accordingly, the present invention provides a pharmaceutical composition which is stable and suitable for oral administration to a patient, comprising a mixture of a substituted pyridylsulfinyl benzimidazole having gastric acid secretion inhibitory activity in an amount sufficient to inhibit gastric acid secretion in said patient, and a pharmaceutically acceptable carrier, said carrier comprising at least one polymer which is at least partially comprised of vinylpyrrolidone monomeric units. Optionally, the mixture also contains other pharmaceutically acceptable excipients. Desirably, the composition is in the form of a simple powder blend or granules of the active ingredient and the carrier, together with any optionally included excipients, filled into an enteric capsule, i.e., a capsule which is coated with an enteric polymer or which is made from an enteric polymer.

The present invention also provides a process for making a pharmaceutical composition which is stable and suitable for oral administration to a patient, comprising mixing together a substituted pyridylsulfinyl benzimidazole having gastric acid secretion inhibitory activity with a pharmaceutically acceptable carrier, the carrier comprising at least one polymer which is at least partially comprised of vinylpyrrolidone monomeric units, together with any optionally included pharmaceutically acceptable excipients. The mixture, which is in the form of a simple powder

blend is then filled into enteric capsules, i.e., capsules which are coated with an enteric polymer

or which are made from an enteric polymer. Alternatively, the mixture, in the form of a powder

blend, is converted into granules and the granules are filled into an enteric capsule.

5 Whereas the processes for preparation of the prior art compositions involve steps for conversion

of a powder blend into core units such as granules, pellets or tablets, and also the step of applying

a subcoat over the core units, the process for preparation of the pharmaceutical composition

according to the present invention does not require these steps because the powder blend or

granules are filled into enteric capsules without being compressed into intact core units, and

10 without applying a subcoat. Furthermore, whereas the core units of prior art compositions contain

several pharmaceutical excipients, the pharmaceutical composition of the present invention may be

a simple mixture comprising a substituted pyridylsulfinyl benzimidazole and one or more

polymers obtained by the polymerization of monomers at least one of which is vinylpyrrolidone.

For the above cited reasons, the process for the preparation of the present invention is simple, less

15 time consuming and more economical than prior art processes.

Detailed Description of the Invention

In accordance with the present invention, the substituted pyridylsulfinyl benzimidazole having

gastric acid secretion inhibitory activity may be one of the following:

(a) a substituted pyridylsulfinyl benzimidazole having the structure of Formula I shown below

wherein R¹ and R² are the same or different, and each of R¹ and R² is selected from the group consisting of hydrogen, halogen, carbomethoxy, carboethoxy, and C₁₋₄ alkyl, alkoxy, or alkanoyl in any position; R⁶ is selected from the group consisting of hydrogen, methyl, and ethyl; R³ and R⁵ are the same or different, and each is selected from the group consisting of hydrogen, methyl, methoxy, ethoxy, methoxyethoxy, and ethoxyethoxy; and R⁴ is methoxy, ethoxy, methoxyethoxy, or ethoxyethoxy; wherein R³, R⁴ and R⁵ are not all hydrogen; OR

15

(b) a substituted pyridylsulfinyl benzimidazole having the structure of Formula II shown below

20

wherein R^1 is hydrogen, methoxy or trifluoromethyl; R^2 and R^3 are independently hydrogen or methyl; R^4 is a $C_{2.5}$ fluorinated alkyl; and n is 0 or 1; OR

(c) a substituted pyridylsulfinyl benzimidazole having the structure of Formula III shown below

wherein R^1 is C_{1-3} alkyl which is at least partially substituted by fluorine, or chlorodifluoromethyl; R^2 is hydrogen, halogen, trifluoromethyl, C_{1-3} alkyl, or C_{1-3} alkoxy which may be partially or completely substituted by fluorine; or R^1 and R^2 together with the oxygen atom to which R^1 is bonded is C_{1-2} alkylenedioxy which may be partially or completely substituted by fluorine, or chlorotrifluoroethylenedioxy; R^4 is C_{1-3} alkoxy; one of R^3 and R^5 is C_{1-3} alkoxy and the other is hydrogen or C_{1-3} alkyl; and n is 0 or 1.

15 Examples of substituted pyridylsulfinyl benzimidazoles that may be used as the active ingredient in the novel compositions of the present invention include omeprazole falling within the definition of Formula I, lansoprazole falling within the definition of Formula II, and pantoprazole falling within the definition of Formula III. The amount of the active agent used in the composition is that which will deliver a suitable therapeutically effective dose, i.e., an amount sufficient to inhibit gastric acid secretion in a patient, on a suitable daily dosing regimen.

According to the present invention, in addition to the substituted pyridylsulfinyl benzimidazole, the pharmaceutical composition contains a carrier comprising one or more polymers that are obtained by polymerization of monomers at least one of which is vinylpyrrolidone.

An example of a class of polymers that may be used in the present invention is polyvinylpyrrolidones also known as povidone or PVP. The United States Pharmacopoeia XXII describes povidone as a synthetic polymer consisting essentially of linear 1-vinyl-2-pyrrolidone groups. The polyvinylpyrrolidones are commonly available from BASF under the brand name Stollidon or from ISP under the brand name Plasdone. Polyvinylpyrrolidone is available as a water soluble polymer or as a cross-linked water insoluble polymer. Examples of water soluble polyvinylpyrrolidones include PVP K-12, PVP K-15, PVP K-17, PVP K-25, PVP K-30, PVP K-60, PVP K-90, and PVP K-120 having approximate molecular weights of 2500, 8000, 10000, 30000, 50000, 400000, 1000000, and 3000000, respectively. Soluble PVP is conventionally used as a binder in tablet formulations. In the present invention, soluble PVP is used in the inventive composition as a stabilizing excipient and as a diluent for the substituted pyridylsulfinyl benzimidazole.

Cross-linked polyvinylpyrrolidone is a polymer obtained by a polymerization process that produces a physically cross-linked polyvinylpyrrolidone (United States Patent No. 3,933,766) which is insoluble in water and in all the usual solvents. Examples of cross-linked polyvinylpyrrolidones that may be used in the present invention include various grades such as those available from BASF under the brand names Kollidon CL, Crospovidone M, and Kollidon CL-M. Because of its high swelling ability, cross-linked polyvinylpyrrolidone is conventionally used as a disintegrant in tablets; however, in the present invention it is used as a stabilizing excipient and as a diluent for the substituted pyridylsulfinyl benzimidazole.

Another example of a class of polymers that may be used in the present invention are water soluble vinylpyrrolidone-vinyl acetate copolymers that are formed by the copolymerization of

vinylpyrrolidone and vinyl acetate. An example of a vinylpyrrolidone-vinyl acetate copolymer that may be used in the present invention is the copolymer available from BASF under the brand name Kollidon VA-64. In the present invention, the vinylpyrrolidone-vinyl acetate copolymer is used as a stabilizing excipient and as a diluent for the substituted pyridylsulfinyl benzimidazole.

5

According to the present invention, the pharmaceutically acceptable carrier is present in an amount from about 10% to about 98%, preferably from about 50% to about 90%, by weight of the total weight of the composition.

According to the present invention, the pharmaceutical composition may also contain conventional pharmaceutically acceptable excipients. Pharmaceutical excipients well known in the pharmaceutical arts can be found listed in the Handbook of Pharmaceutical Excipients (Ed. A. Wade and P.J. Weller, The Pharmaceutical Press, London), in the U.S. FDA listing of inactive ingredients, and in other sources of pharmaceutical literature.

15

In preferred embodiments, the pharmaceutically acceptable excipients may comprise fatty acid glycerides. One example of fatty acid glycerides that may be used in the invention is a mixture of glycerides (e.g., mono-, di- and/or triglycerides) of long chain (e.g., C₁₂ - C₁₈) fatty acids; for example, the range of products available under the brand name Gelucire (Gattefosse Corporation).

20 Another example of fatty acid glycerides that may be used in the invention is a mixture of glycerides (e.g., triglycerides) of medium chain length (e.g., C₈ - C₁₀) fatty acids; for example, the range of products available under the brand names Miglyol, Crodamol GTC/C, MCT oil, Neobee M5, AKOMED, Nesatol, and the like. The fatty acid glycerides included in the composition of this invention can also be in the form of vegetable oils, such as castor oil,

hydrogenated castor oil, or hydrogenated vegetable glycerides, such as those available under the brand name Witepsol.

According to the process of the present invention, a substituted pyridylsulfinyl benzimidazole

5 having gastric acid secretion inhibitory activity, a carrier comprising one or more polymers
comprising vinylpyrrolidone monomeric units, optionally together with pharmaceutically
acceptable excipients, are mixed together to obtain a blend or granules, the blend or granules so
obtained are filled into capsules, and the capsules are then enteric coated, or the blend or granules
are filled into capsules having an enteric coating or made from an enteric material. In

10 embodiments where one of the pharmaceutically acceptable excipients is a fatty acid glyceride, a
liquid fatty acid glyceride is mixed with the other ingredients of the composition, or a solid fatty
acid glyceride is first heated to above its melting point and the liquid obtained mixed with other
ingredients of the composition to obtain granules.

The capsules used in the invention may be hard or soft capsules. The outer shell of the capsules may be composed of a film forming agent or agents, water and plasticizer. The shell may also contain coloring and opacifying agents. Examples of film forming agents which may be used in the capsule shell include gelatin, methylcellulose, hydroxypropyl methylcellulose, hydroxypropyl cellulose, and the like. When the shell is of a conventional type, e.g., it is made from gelatin, it is given an outer enteric coat. Alternatively, the capsules are enteric capsules wherein the shell itself is enteric in nature. The shell of enteric capsules may be made from one or more film forming polymers at least one of which has an enteric nature. The composition of enteric capsules is a known art. For example, the shell may be made from a mixture of polymers such as gelatin or hydroxypropyl methylcellulose, and one or more enteric polymers, such as a polyacrylate

enteric polymer, cellulose acetate phthalate, hydroxypropyl methyl cellulose acetate succinate, or cellulose acetate butyrate; or from a mixture of gelatin or hydroxypropyl methylcellulose and polyvinyl acetate phthalate; or from calcium alginate and the like. The enteric polymers may be present as free acids or their salts. When the shell is of a conventional type, an outer enteric coating may be applied using known art. The coating composition may be aqueous or organic solvent based. The drying of the applied layers of a coating composition may be achieved by conventional means or by application of vacuum.

In another embodiment of the process of the present invention, a substituted pyridylsulfinyl benzimidazole having gastric acid secretion inhibitory activity, a carrier comprising one or more vinylpyrrolidone polymers, optionally together with pharmaceutically acceptable excipients, are mixed together to obtain a powder blend, and the blend so obtained is subjected to conventional processing steps to obtain granules or tablets.

15 The present invention is further illustrated by the following non-limiting examples.

EXAMPLE 1

Omeprazole and cross-linked polyvinylpyrrolidone in amounts as given in Table 1 were mixed together. The blend so obtained was filled into capsules.

TABLE 1

	Ingredient	Weight (mg/capsule)
	Omeprazole	20.00
5	Cross-linked polyvinylpyrrolidone (Kollidon CL-M)	100.00
	Total	120.00

10

The capsules were enteric coated in a Freund Hi-coater to a weight build-up of 10% using the coating composition as given in Table 2.

TABLE 2

15

Ingredient	Weight (g)
Eudragit L - 100 - 55	100.00
Sodium hydroxide	1.40
Titanium dioxide	170
Talc	50.00
Polyethylene glycol-300	20.00
Water	650.00

25

20

EXAMPLE 2

Omeprazole and vinylpyrrolidone-vinyl acetate copolymer in amounts as given in Table 3 were mixed together. The blend so obtained was filled into capsules.

TABLE 3

	Ingredient	Weight (mg/capsule)
	Omeprazole	20.00
5	Vinylpyrrolidone-vinyl acetate copolymer (Kollidon VA-64)	100.00
	Total	120.00

The capsules were enteric coated in a Freund Hi-coater to a weight build-up of 10% using the coating composition as given in Table 2.

EXAMPLE 3

Omeprazole, vinylpyrrolidone-vinyl acetate copolymer, and cross-linked polyvinylpyrrolidone in amounts as given in Table 4 were mixed together. The blend was filled into capsules.

TABLE 4

Ingredient	Weight (mg/capsule)
Omeprazole	20.00
Cross-linked polyvinylpyrrolidone (Kollidon CL-M)	50.00
Vinylpyrrolidone-vinyl acetate copolymer (Kollidon VA-64)	50.00
Total	120.00

25

20

The capsules were enteric coated in a Freund Hi-coater to a weight build-up of 10% using the coating composition as given in Table 2.

EXAMPLE 4

Omeprazole and polyvinylpyrrolidone (PVP K 30) in the amounts as given in Table 5 were mixed together. The blend so obtained was filled into capsules.

5

TABLE 5

Ingredient	Weight (mg/capsule)
Omeprazole	20.00
PVP K30	100.00
Total	120.00

10

The capsules were enteric coated in a Freund Hi-coater to a weight build-up of 10% using the coating composition as given in Table 2.

15

EXAMPLE 5

Omeprazole and the other ingredients in the amount as given in Table 6 were mixed together. The blend so obtained was filled into capsules.

20

TABLE 6

Ingredient	Weight (mg/capsule)
Omeprazole	20.00
Kollidon CL-M	50.00
Avicel PH 112	50.00
Total	120.00

25

The capsules were enteric coated in a Freund Hi-coater to a weight build-up of 10%.

EXAMPLE 6

Omeprazole and Kollidon CL-M in amounts as given in Table 7 were mixed together. AKOMED R (fatty acid glyceride composed of caprylic / capric triglycerides and derived from coconut and/or palm kernel oils) and Gelucire 33/01 (a mixture of glycerides, e.g., mono-, di- and/or triglycerides of long chain fatty acids) were heated to 60°C for 20 minutes, stirred well and cooled to 30°C. The omeprazole and the Kollidon blend was granulated with the liquid mix. The granules were screened through a No. 22 mesh sieve and filled into capsules. The capsules were enteric coated in a Freund Hi-coater to a weight build-up of 10% using the coating composition as given in Table 2.

10

TABLE 7

Ingredient	Weight (mg/capsule)
Omeprazole	20.00
Cross-linked polyvinylpyrrolidone (Kollidon CL-M)	100.00
Gelucire 33/01	10.00
AKOMED R	20.00
Total	150.00

20

15

The enteric coated capsules of Examples 1 to 6 were tested as described under dissolution test (Method B) for delayed release (enteric coated) dosage forms in the United States Pharmacopoeia XXIII, page 1795. In the acid stage, omeprazole was not released from the capsules. The data for percent released in the buffer stage is given in Table 8.

25

TABLE 8

ſ	TIME (MINUTES)	I	MEAN CUI	MULATIV	E PERCEN	T RELEASEI	EASED			
	` ,	Ex. 1	Ex. 2	Ex. 3	Ex. 4	Ex. 5	Ex. 6			
	20	6.50	21.60	58.40	14.70	1.70	72.20			
	30	46.70	39.50	83.66	30.00	4.80	102.00			
T	45	91.50	60.10	95.60	75.00	84.90	106.80			

In another test, the enteric coated capsules of Examples 1 to 6 were kept in high density polyethylene bottles at 40°C/ 75% RH (Relative Humidity). The pharmaceutical compositions filled in the enteric capsules did not show any sign of instability such as change in color or appearance as given in Table 9.

5

TABLE 9

	OBSERVATION				
EXAMPLE No.	15 days, 40°C/75% RH	30 days, 40°C/75% RH			
1	No Change	No Change			
2	No Change	No Change			
3.	No Change	No Change			
4	No Change	No Change			
5	No Change	No Change			
6	No Change	No Change			

15

10

The Omeprazole content in the capsules stored as given above for a period of 30 days was determined by a stability-indicating HPLC method. The results are given in Table 10.

TABLE 10

20

EXAMPLE No.	Assay
1	105.10
2	100.19
3	99.85
4	99.66
5	98.07
6	95.20

25

While the invention has been described by reference to specific embodiments, this was for purposes of illustration only. Numerous alternative embodiments will be apparent to those skilled in the art and are considered to be within the scope of the claimed invention.

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CLAIMS

- 1. A pharmaceutical composition which is stable and suitable for oral administration to a patient, comprising a mixture of a substituted pyridylsulfinyl benzimidazole having gastric acid secretion inhibitory activity in an amount sufficient to inhibit gastric acid secretion in said patient, and a pharmaceutically acceptable carrier, said carrier comprising at least one polymer having vinylpyrrolidone monomeric units.
- 2. The composition of claim 1 wherein the substituted pyridylsulfinyl benzimidazole has the structure of Formula I

$$R^2$$
 N
 S
 CH
 N
 R^4
 R^5
 R^5
 R^6
 R^6

wherein R¹ and R² are the same or different, and each of R¹ and R² is selected from the group consisting of hydrogen, halogen, carbomethoxy, carboethoxy, and C₁₋₄ alkyl, alkoxy, or alkanoyl in any position; R⁶ is selected from the group consisting of hydrogen, methyl, and ethyl; R³ and R⁵ are the same or different, and each is selected from the group consisting of hydrogen, methyl, methoxy, ethoxy, methoxyethoxy, and ethoxyethoxy; and R⁴ is methoxy, ethoxy, methoxyethoxy or ethoxyethoxy; wherein R³, R⁴ and R⁵ are not all hydrogen.

3. The composition of claim 1 wherein the substituted pyridylsulfinyl benzimidazole has the structure of Formula II

wherein R^1 is hydrogen, methoxy or trifluoromethyl; R^2 and R^3 are independently hydrogen or methyl; R^4 is a $C_{2.5}$ fluorinated alkyl; and n is 0 or 1.

4. The composition of claim 1 wherein the substituted benzimidazole has the structure of Formula III

$$R^2$$
 N
 S
 CH_2
 N
 R^5
 R^5
 R^5
 R^1
 R^1
 R^0
 R^0

10

15

5

wherein R^1 is $C_{1.3}$ alkyl which is at least partially substituted by fluorine, or chlorodifluoromethyl; R^2 is hydrogen, halogen, trifluoromethyl, $C_{1.3}$ alkyl, or $C_{1.3}$ alkoxy which may be partially or completely substituted by fluorine; or R^1 and R^2 together with the oxygen atom to which R^1 is bonded is $C_{1.2}$ alkenedioxy which may be partially or completely substituted by fluorine, or chlorotrifluoroethylenedioxy; R^4 is $C_{1.3}$ alkoxy; one of R^3 and R^5 is $C_{1.3}$ alkoxy and the other is hydrogen or $C_{1.3}$ alkyl; and n is 0 or 1.

The composition of claim 1 wherein the substituted benzimidazole

The composition of claim 1 wherein said polymer is polyvinyl pyrrolidone.

20

5. The composition of claim 1 wherein the substituted benzimidazole comprises omegrazole.

comprises pantoprazole.

6.

8.

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7. The composition of claim 1 wherein the substituted benzimidazole comprises lansoprazole.

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9. The composition of claim 1 wherein said polymer is cross-linked polyvinylpyrrolidone.

PCT/IB99/00139 WO 99/61022

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	10. The composition of claim 1 wherein said polymer is a vinylpyrrolidone-
	vinyl acetate copolymer.
	11. The composition of claim 1 wherein said carrier comprises about 10% to
5	about 98% by weight of said mixture.
	12. The composition of claim 1 wherein said carrier comprises about 50% to
	about 90% by weight of said mixture.
10	13. The composition of claim 1 wherein said mixture further comprises a fatty
10	acid glyceride.
	acia processes.
	14. The composition of claim 1 in the form of a capsule, said mixture being
	contained within a capsule shell made from an enteric material.
15	
	15. The composition of claim 14 wherein said mixture contained within said
	capsule shell is in the form of a powder blend.
	16. The composition of claim 14 wherein said mixture contained within said
20	capsule shell is in the form of granules.
	17. The composition of claim 1 in the form of a capsule, said mixture being
	contained within a capsule shell which is coated with an enteric material.
	Contained within a capable circle was a capable with a capable circle was an extensive and
25	18. The composition of claim 17 wherein said mixture contained within said
	capsule shell is in the form of a powder blend.
	19. The composition of claim 17 wherein said mixture contained within said
	capsule shell is in the form of granules.
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	20. The composition of claim 1 in the form of a tablet.

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- 21. A process for preparing a stable pharmaceutical composition which is suitable for oral administration, comprising mixing a substituted pyridyl sulfinyl benzimidazole having gastric acid secretion inhibitory activity in an amount sufficient to inhibit gastric acid secretion in a patient together with and a pharmaceutically acceptable carrier, said carrier comprising at least one polymer having vinylpyrrolidone monomeric units to form a mixture.
- 22. The process of claim 21 wherein the substituted pyridyl sulfinyl benzimidazole has the structure of Formula I

$$R^2$$
 R^3
 R^4
 R^5
 R^1
 R^4
 R^5
 R^5
 R^6
 R^6

wherein R¹ and R² are the same or different, and each of R¹ and R² is selected from the group consisting of hydrogen, halogen, carbomethoxy, carboethoxy, and C_{1.4} alkyl, alkoxy, or alkanoyl in any position; R⁶ is selected from the group consisting of hydrogen, methyl, and ethyl; R³ and R⁵ are the same or different, and each is selected from the group consisting of hydrogen, methyl, methoxy, ethoxy, methoxyethoxy, and ethoxyethoxy; and R⁴ is methoxy, ethoxy, methoxyethoxy or ethoxyethoxy; wherein R³, R⁴ and R⁵ are not all hydrogen.

23. The process of claim 21 wherein the substituted pyridyl sulfinyl benzimidazole has the structure of Formula II

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wherein R^1 is hydrogen, methoxy or trifluoromethyl; R^2 and R^3 are independently hydrogen or methyl; R^4 is a $C_{2.5}$ fluorinated alkyl; and n is 0 or 1.

24. The process of claim 21 wherein the substituted benzimidazole has the structure of Formula III

wherein R¹ is C_{1.3} alkyl which is at least partially substituted by fluorine, or chlorodifluoromethyl; R² is hydrogen, halogen, trifluoromethyl, C_{1.3} alkyl, or C_{1.3} alkoxy which may be partially or completely substituted by fluorine; or R¹ and R² together with the oxygen atom to which R¹ is bonded is C_{1.2} alkenedioxy which may be partially or completely substituted by fluorine, or chlorotrifluoroethylenedioxy; R⁴ is C_{1.3} alkoxy; one of R³ and R⁵ is C_{1.3} alkoxy and the other is hydrogen or C_{1.3} alkyl; and n is 0 or 1.

- 25. The process of claim 21 wherein the substituted benzimidazole comprises omeprazole.
- 26. The process of claim 21 wherein the substituted benzimidazole comprises pantaprazole.
 - 27. The process of claim 21 wherein the substituted benzimidazole comprises lansoprazole.
 - 28. The process of claim 21 wherein said polymer is polyvinylpyrrolidone.

29.	The process of claim 21 wherein said polymer is cross-linked
polyvinylpyrro	lidone.

- 30. The process of claim 21 wherein said polymer is a vinylpyrrolidine-vinyl acetate copolymer.
 - 31. The process of claim 21 further comprising mixing at least one pharmaceutically acceptable excipient into said mixture.
- 10 32. The process of claim 31 wherein said pharmaceutically acceptable excipient is a fatty acid glyceride.
 - 33. The process of claim 21 further comprising filling said mixture in the form of a powder blend into a capsule made from an enteric material.
 - 34. The process of claim 21 further comprising coating a capsule shell with an enteric material, and filling said mixture in the form of a powder blend into said capsule shell.
 - 35. The process of claim 21 further comprising granulating said mixture to produce granules, and filling said granules into a capsule shell made from an enteric material.
 - 36. The process of claim 21 further comprising granulating said mixture so as to form granules, coating a capsule shell with an enteric material, and filling said granules into said capsule shell.
 - 37. The process of claim 21 further comprising compressing said mixture into tablets.

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int itional Application No PCT/IB 99/00139

. classification of subject matter PC 6 A61K31/44 A61K A61K9/48 A61K47/32 According to International Patent Classification (IPC) or to both national classification and IPC **B. FIELDS SEARCHED** Minimum documentation searched (classification system followed by classification symbols) IPC 6 Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched Electronic data base consulted during the international search (name of data base and, where practical, search terms used) C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Citation of document, with indication, where appropriate, of the relevant passages Category ° WO 97 12580 A (PHARMA PASS LLC ; SETH PAWAN 1-37 X (US)) 10 April 1997 cited in the application * p.3, 1.24-25; p.7, 1.22-39; p.8, 1.35; p.10, 1.12-13; p.12, 1.19; Ex. 1-5 and 15-20; claims 8-10 *1 - 37WO 96 01623 A (ASTRA AB ; BERGSTRAND PONTUS Х JOHN ARVID (SE); LOEVGREN KURT INGMAR) 25 January 1996 * p.6, 1.24; p.8, 1.3, 1.11 and 1.16; Ex. 8, 16 and 17 * US 5 178 867 A (GUITTARD GEORGE V ET AL) 1 - 37χ 12 January 1993 * col.6, 1.43; col.5, 1.41-55; col.9, 1.4; Ex.4; claim 10 * -/--Patent family members are listed in annex. Further documents are listed in the continuation of box C. Χ ° Special categories of cited documents : "T" later document published after the international filing date or priority date and not in conflict with the application but "A" document defining the general state of the art which is not considered to be of particular relevance cited to understand the principle or theory underlying the invention "E" earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to filing date "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) involve an inventive step when the document is taken alone "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-"O" document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled in the art. document published prior to the international filing date but later than the priority date claimed "&" document member of the same patent family Date of the actual completion of the international search Date of mailing of the international search report 21/05/1999 6 May 1999 Authorized officer Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentlaan 2 NL - 2260 HV Rijswijk Tel. (+31-70) 340-2040, Tx. 31 651 epo nl, Uiber, P Fax: (+31-70) 340-3016

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